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**Natural history of epithelial ovarian cancer and its relation to surgical and medical treatment**

**De Iaco P *et al*.** EOC surgical and medical treatment

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**Abstract**

Epithelial ovarian cancer (EOC) represents approximately 90% of primary malignant ovarian tumors, the sixth most common cancer in women and the second most common gynecologic cancer. Approximately 80%–85% of all ovarian carcinomas in Western society are serous and up to 95% of patients are in advanced stages (FIGO stage III–IV) at diagnosis. Treatment of ovarian cancer is mainly based on three key approaches: surgical removal of neoplasia; chemotherapy to kill cancer cells; direct chemotherapy on peritoneal surfaces. The application of hyperthermic chemotherapy to the peritoneal cavity (HIPEC) after radical surgery may also be an attractive option. We analyzed the natural history of EOC in the literature and identified various time-points where sensitivity to chemotherapy, freedom from disease and overall survival are different. We propose eight time-points in EOC history with homogeneous oncological findings. The effectiveness of HIPEC in EOC treatment should be evaluated based on these eight time-points and we believe that retrospective and prospective studies of HIPEC should be evaluated according to these time-points.

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**Key words:** Ovarian cancer; HIPEC; Chemo-sensitivity; Time-points; Survival

**Core tip:** The standard treatment for advanced ovarian cancer consists in complete cytoreductive surgery and intravenous combination chemotherapy with a platinum compound and a taxane. Although response rates to initial therapy are high, many patients will recur and die of peritoneal carcinomatosis. The addition of hyperthermic intraperitoneal chemotherapy (HIPEC) to the standard therapy aims at increasing survival by reducing peritoneal recurrence. In this review we discuss the time points where HIPEC can be proposed.

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**INTRODUCTION**

Epithelial ovarian cancer (EOC) represents approximately 90% of primary malignant ovarian tumors, the sixth most common cancer in women and the second most common gynecologic cancer[1]. At diagnosis, the majority of patients (70%) are in advanced stage of the disease (FIGO IIB-IV) with rapid and asymptomatic widespread cancer cells in the pelvic structure and peritoneal cavity[2]. Disease stage at presentation is the most important prognostic factor determining outcome; 70%–80% of women at stage I survive for five years compared to only 15% of those at stage IV[3].

Recent advances in pathology and genetics have shown that EOC is a heterogeneous disease with various risk factors, genetic abnormalities and oncological pathways that partly determine biological behavior, response to chemotherapy, and prognosis[4]. A dualistic model places the major histological types into two groups: types I and II. Type I cancers (mucinous, endometrioid, clear cell carcinomas and low-grade serous carcinomas) demonstrate a relatively insidious clinical course with generally better prognosis. These develop in a stepwise fashion from well-established precursor lesions, such as borderline tumors and endometriosis[5,6]. Type I are relatively genetically stable and typically display a variety of somatic mutations in genes including *K-ras*, *BRAF*, *PTEN*, *CTNNB1* but very rarely TP53. In contrast, Type II cancers (high-grade serous carcinomas, high-grade transitional carcinomas, malignant mixed mesodermal tumors and undifferentiated carcinomas) are extremely aggressive neoplasms with remarkable early sensitivity to platinum-based chemotherapy but are frequently diagnosed at advanced stages. They are chromosomally highly unstable and harbour TP53 mutations in more than 95% of cases[7,8]. Approximately 80%–85% of all ovarian carcinomas in Western society are serous. Up to 95% of patients with FIGO stage III–IV disease have serous carcinomas whereas FIGO stage I serous carcinomas are uncommon[9,10].

Like other cancers, EOC can spread through lymphatic and blood vessels to nodes and parenchyma of distant organs (liver, lung and brain). However, a distinctive feature of these tumors is their ability to spread from the ovary to the abdominal cavity, forming nodules of variable size on the surface of the parietal and visceral peritoneum, including the omentum. The coalescence of nodules forms plaque or masses in the abdominal-pelvic cavity. Blockage of diaphragmatic lymphatics prevents outflow of proteinaceous fluid from the peritoneal cavity, causing the accumulation of ascites in advanced disease. Tumor dissemination from the peritoneal cavity to the pleural cavity occurs through the diaphragmatic peritoneum and leads to pleural effusion[11,12].

**TREATMENT OF PRIMARY ADVANCED OVARIAN CANCER**

The main treatment of advanced disease consists of surgical removal of all visible nodules in the abdominal cavity followed by intravenous chemotherapy (platinum-based drugs with or without taxanes). The combined effect of surgery and chemotherapy is often the complete eradication of cancer cells.

Treatment of ovarian cancer is mainly based on three key areas: surgical removal of neoplasia; chemotherapy to kill cancer cells; application of chemotherapy directly on peritoneal surfaces.

In advanced disease (FIGO stage IIB-IIIC) the surgical removal of neoplasia with optimal cytoreduction (nodules ≤ 1 cm left) is recommended. An additional survival advantage of complete cytoreduction (no visible residual disease) has been recently reported[12]. Several studies have shown that specialized gynecological oncologist surgeons are more likely to perform optimal surgery than general surgeons[13]. The frequent presence of multiple neoplastic implants on peritoneal surfaces together with pelvic and upper abdominal organs implies that surgeons must be prepared to remove organs beyond the pelvis, such as peritoneal surfaces of colic gutters, diaphragmatic domes, and to carry out surgical procedures on the colon, bowel, liver, gallbladder, stomach, and spleen. This implies multidisciplinary surgical effort and the possibility of higher postoperative morbidity. This idea has not been accepted by the majority of gynecologic oncologists due to the lack of scientific data. If initial maximal cytoreduction is not carried out, interval debulking surgery (IDS) should be considered in patients responding to chemotherapy or with stable disease. IDS should ideally be carried out after three cycles of chemotherapy then followed by three further chemotherapy cycles[14].

Chemotherapeutic efficacy for ovarian carcinoma showed a dramatic shift after the introduction of platinum compounds and since 1996 the combination of platinum plus paclitaxel has been the standard treatment. The current rationale of six cycles of treatment as standard is based on three randomized trials which analyzed the impact of chemotherapy duration (*i.e.*, number of cycles) on OS. None of these studies demonstrated a difference in median survival time, but longer durations were associated with increased toxicity, especially neuropathy[15]. Other chemotherapy regimens, such as gemcitabine and liposomal doxorubicin in association with carboplatin–paclitaxels were compared to carboplatin-plaxitel alone in the Phase III Gynecologic Cancer Intergroup (GCIG) trials (GOG 0182-ICON 5). These showed no statistically significant superiority or clinically useful benefit associated with the three drugs compared to the controls. Currently carboplatin–paclitaxel remains the treatment of choice even though angiogenesis inhibitors in combination with the standard treatment have been approved by the US Food and Drug Administration[16]. The main issue with EOC is the chemo sensitivity of cancer cells. Data shows that only 50% of patients have a complete clinical response to standard IV chemotherapy and that 30% of them have microscopic metastasis at surgical exploration. Most advanced stage patients who achieve clinical remission after completion of initial treatment develop recurrent disease and drug resistance, and their cure rate is less than 30%. These factors are major limitations in the treatment of patients with EOC. In order to overcome these limitations, different treatments such as secondary cytoreduction, second line chemotherapy drugs, high dose chemotherapy, intra-peritoneal chemotherapy (IP), radiotherapy, immunotherapy and hormone therapy should be considered. To date, none of these approaches, apart from IP chemotherapy, has been found to have a significant impact on survival[17].

IP chemotherapy refers to the administration of cytotoxic agents directly to the peritoneal cavity. The rationale is that a higher concentration of cytotoxic drugs and longer duration of exposure can be achieved while reducing the toxicity normally associated with intravenous therapy[18-20]. In fact, IP administered cytotoxic drugs can directly target tumor masses confined to the abdominal cavity, thus bypassing the poor vascularization of small volume disease and thereby increasing peri- and intra-tumoral drug concentration. Cisplatin can penetrate small volume tumors to a maximum depth of 1-3 mm and may therefore only benefit those patients with microscopic residual disease. By using large intra-peritoneal doses, the tumor surface can be exposed to high concentrations of cisplatin with only a small amount of drug leaking into the circulation. By this means, the amount of cisplatin reaching the tumor through capillaries is doubled when compared to the maximum tolerated dose delivered intravenously. Several studies have documented the advantages of IP compared to IV chemotherapy[20]. Post-operative adhesions after cytoreductive surgery can limit the access of the active drug to tumor areas and other complications, such as infections due to the IP catheter, may occur. Intra-operative administration of IP chemotherapy has been designed to overcome such obstacles. Intra-peritoneal hyperthermia chemotherapy (HIPEC) is a new treatment method based on increasing the sensitivity of cancer cells to the direct cytotoxic effect of chemotherapeutic agents at high temperature and increasing the concentration of chemotherapeutic agents that penetrate cancer tissues [21-23].

**TREATMENT OF RECURRENCE**

Approximately 70% of patients with advanced cancer who experience clinical remission after initial surgery and chemotherapy will develop recurrent disease [24].

In general, patients who progress during treatment with platinum are considered to have ‘platinum-refractory’ disease and patients who show recurrence < 6 mo after completion of first-line platinum chemotherapy are considered to have ‘platinum resistant’ disease. These patients are candidates for salvage therapy with second line chemotherapy. Patients who relapse after an interval of > 6-12 mo are defined as ‘platinum-sensitive’ and are candidates for chemotherapy and/or surgery. The concept of chemo-sensivity is based on clinical data; re-subjecting the patient to the previous chemotherapy regimen obtains about a 20% response, but drug administration is the only method by which to verify cell response. Because of the late onset of relapse, platinum-sensitive patients should in reality be regarded as including both chemo-sensitive and chemo-insensitive patients[25].

Appropriate treatment of recurrence (chemotherapy/surgery), which may be based on time and nature of relapse and the role of surgery, remains a field of discussion and controversy.

In general, surgical resection may be considered in platinum-sensitive patients. Resectable disease, good performance status and complete secondary cytoreduction are one of the best predictors of survival in these patients[26-28].

In ovarian platinum-sensitive recurrence, surgical cytoreduction offers the following potential benefits: (1) cytoreduction of tumor volume offers patients a greater chance of response to chemotherapy; and (2) the elimination of potentially chemo-resistant cells. However, surgical cytoreduction is generally not undertaken without also scheduling postoperative chemotherapy since surgery alone rarely offers a cure.

**ADVANCED EOC NATURAL HISTORY TIME-POINTS**

We analyzed literature using the search terms “ovarian cancer” and “HIPEC treatment”. EOC naturally presents various time-points where surgery, chemotherapy or HIPEC can be identified with homogenous chemo-sensitivity, response to therapy, and survival. Chua *et al*[27] proposed five time-points: (1) time of primary treatment; (2) time of IDS; (3) time of consolidation therapy after complete pathological response following initial therapy; (4) time of first recurrence; or (5) time of salvage therapy (Figure 1). The results of the most important paper are shown in table 1 [28,29].

Given that chemo-sensitivity is an important issue for the prognosis and the homogeneity of these patients, we considered eight time-points upon which a clinical trial could be based: (1) time of primary treatment where optimal cytoreduction is achieved (group with chemo-sensitive and chemo-insensitive tumors), (2)time of IDS after neo-adjuvant chemotherapy with partial or complete response (chemo-sensitive group), (3)time of IDS after neo-adjuvant chemotherapy with stable disease (chemo-insensitive group), (4) time of consolidation therapy after complete pathological response following initial therapy (chemo-sensitive group), (5)time of first recurrence when disease relapses more than 12 mo after treatment (chemo-sensitive/chemo-insensitive group), (6)time of first recurrence when disease relapses more than 12 mo after treatment and a course of chemotherapy obtains complete response (chemo-sensitive group), (7) time of first recurrence when disease relapses less than 12 mo after treatment (chemo-insensitive group) (8) time of salvage therapy after various chemotherapy lines (chemo-insensitive group) (Figure 2). Correct analysis of past and future clinical trials should take account of these time-points in patient evaluation.

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**Figure 1 Epithelial ovarian cancer natural history: time-points where** **intra-peritoneal hyperthermia chemotherapy can be proposed.**

**Figure 2 Epithelial ovarian cancer natural history: time-points where intra-peritoneal hyperthermia chemotherapy can be proposed and where chemo-sensitivity and chemo-insensitivity were evaluated.**

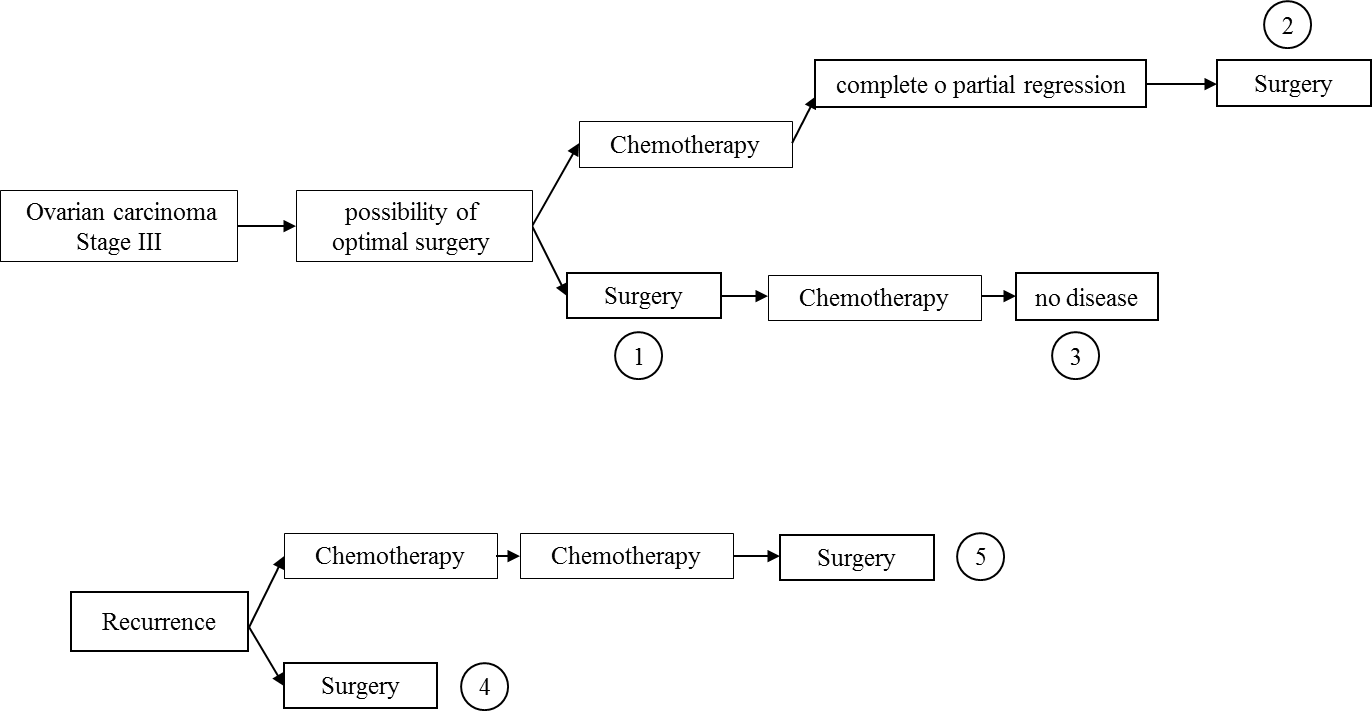


Fig 1

Figures in the circles represent time-points of natural history of EOC where Hipec can be proposed (Chua 2009 modified)



Fig 2

Figures in the circles represent time-points of natural history of EOC where Hipec can be proposed where chemosensitivity and chemoinsensitivity were evaluated (see the text).

Table 1

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | Patients n | Time-point of optimal cytoreduction | Median disease free survival (months) | Overal 3-years survival (%) | Overal 5-years survival (%) |
| Ansaloni et al. (2012) | 39 | 1, 2, 3, 4, 5 | 42\* | NR | NR |
| Deraco et al. (2012) | 56 | 4, 5 | NR | NR | 23 |
| Pomel et al. (2010) | 31 | 2, 3 | 27 | 67° | NR |
| Bereder et al. (2009) | 246 | 2, 4, 5 | 13 | 60 | 35 |
| Pavlov et al. (2009) | 56 | 1, 4, 5 | 26 | NR | NR |
| Fagotti et al. (2009) | 25 | 4, 5 | 10 | NR | NR |
| Guardiola et al. (2009) | 47 | 2 | 14 | 63° | NR |
| Di Giorgio et al. (2008) | 47 | 1, 4, 5 | 20 | NR | 17 |
| Bae et al. (2007) | 67 | 2, 3 | NR | NR | 66 |
| Cotte et al. (2007) | 81 | 5 | 19 | NR | NR |
| Helm et al. (2007) | 18 | 5 | 10 | NR | NR |
| Rufian et al. (2006) | 33 | 1, 4 | NR | 46 | 37 |
| Raspagliesi et al. (2006) | 40 | 3, 5 | 11 | NR | 15 |
| Reichman et al. (2005) | 13 | 1, 4 | 15 | 55 | NR |
| Gori et al. (2005) | 29 | 3 | 57\* | NR | NR |
| Look et al. (2004) | 28 | 1, 5 | 17 | NR | NR |
| Ryu et al. (2004) | 57 | 2, 3 | 26 | NR | 54 |
| Piso et.al (2004) | 19 | 1, 4, 5 | 18 | NR | 15 |
| Zanon et al (2004) | 30 | 2 | 17 | 35 | 12 |
| Chatzigeorgiou et al. (2003) | 20 | 5 | 21 | NR | NR |
| de Bree et al. (2003) | 19 | 4, 5 | 26 | 63 | 42 |
| Cavaliere et al (2000) | 20 | NR | NR | 50° | NR |

Table 2: Time-points of optimal cytoreduction and survival result of cytoreductive surgery and hypertermic intraperitoneal chemotherapy of 1021 patients with carcinomatosis from ovarian cancer.

\* refers to results expressed as mean

° 2-years survival result

|  |
| --- |
|  |